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2019-08

This is the pre-peer reviewed version of the following article:

Price SJ, Leung WTM, Owen CJ, et al. Effects of historic and projected climate change on the range and impacts of an emerging wildlife disease. *Glob Change Biol.* 2019;00:1–13

which has been published in final form at <https://doi.org/10.1111/gcb.14651>.

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Effects of historic and projected climate change on the range and impacts of an emerging wildlife disease

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Paper type: Primary Research Article

Running Title: Climate change impacts on a wildlife epidemic

Abstract

The global trend of increasing environmental temperatures is often predicted to result in more severe disease epidemics. However, unambiguous evidence that temperature is a driver of epidemics is largely lacking, because it is demanding to demonstrate its role among the complex interactions between hosts, pathogens and their shared environment. Here we apply a three-pronged approach to understand the effects of temperature on ranavirus epidemics in common frogs, combining *in vitro*, *in vivo* and field studies. Each approach suggests that higher temperatures drive increasing severity of epidemics. In wild populations, ranavirosis incidents were more frequent and more severe at higher temperatures, and their frequency increased through a period of historic warming in the 1990s. Laboratory experiments using cell culture and whole animal models showed that higher temperature increased ranavirus propagation, disease incidence, and mortality rate. These results, combined with climate projections, predict severe ranavirosis outbreaks will occur over wider areas and an extended season, affecting larval recruitment. Since ranaviruses affect a variety of ectothermic hosts (amphibians, reptiles and fish), wider ecological damage is likely. Our three complementary lines of evidence present a clear case for direct environmental modulation of these epidemics and suggest management options to protect species from disease.

Keywords: ranavirus, temperature, virulence, climate change, emerging infectious disease, host-pathogen interactions, common frog, *Rana temporaria*, amphibian population decline

1 Introduction

Interactions between hosts, pathogens and their shared environment shape infectious disease outcomes, the timing of outbreaks, and the invasiveness of pathogens (Engering, Hogerwerf, & Slingenbergh, 2013). Climatic conditions at local and landscape scales represent a critical dimension of the host environment - affecting behavior (e.g. aggregation), stress and immunity - but can also directly affect pathogen (and vector) growth and survival (Altizer, Ostfeld, Johnson, Kutz, & Harvell, 2013; A. Dobson, Molnár, & Kutz, 2015; Engering et al., 2013; Epstein, 2001; Grassly & Fraser, 2006). As such, climatic conditions modulate host-pathogen interactions and operate on multiple timescales: acting annually in driving seasonality (Altizer et al., 2006; Grassly & Fraser, 2006) as well as over longer time periods in determining responses to climate change (Dobson et al., 2015) and influencing the rate and pattern of invasions by emerging pathogens (e.g. Seimon et al., 2007).

Despite this, invasion through pathogen range expansion and environmental change have frequently been viewed as mutually exclusive factors in explaining disease emergence, as indicated clearly in the framing and use of the “novel pathogen” (spread of an exotic pathogen through naïve populations) and “endemic pathogen” (emergence due to perturbations of interaction between hosts and native pathogens) hypotheses (Rachowicz et al., 2005). The “global panzootic” in amphibians caused by the fungal pathogen, *Batrachochytrium dendrobatidis*, serves as a prominent example of how climate change and pathogen range expansion have been pitted against one another as alternative explanations of declines (Berger et al., 1998; Lips, Diffendorfer, Mendelson III, & Sears, 2008; Pounds et al., 2006). There is strong evidence supporting the rapid international spread of a global panzootic lineage of *B. dendrobatidis* during the 20th century (Farrer et al., 2011; O’Hanlon et al., 2018), but the proposed conflict between the two hypotheses now appears to have been reconciled in a framework that incorporates both as key drivers of emergence and outcomes, explaining observations of decline in regions where the impacts have been greatest (Cohen, Civitello, Venesky, McMahon, & Rohr, 2018; Cohen et al., 2017; Raffel et al., 2013). Thus, it seems likely that the previous mindset of treating environmental change and pathogen range expansion as conflicting has hampered understanding of the patterns of emergence and the focusing of mitigation efforts.

Establishing a role for climate in disease emergence can be very challenging. Increasing environmental temperature is a key component of climate change, which is cited as a driver of infectious disease emergence and severity, but evidence for this is scarce and it is often difficult to discriminate between the effect of temperature and other aspects of climate (Harvell et al., 2002). The direct and indirect influences of temperature on host-pathogen interactions (Altizer et al., 2006; Clare et al., 2016; Garner, Rowcliffe, & Fisher, 2011) and its nonlinear effects on incidence and severity (Bosch, Carrascal, Durán, Walker, & Fisher, 2007; Raffel et al., 2013; Walker et al., 2010) represent considerable challenges to a better understanding of disease emergence. Most research effort in this area has focused on human diseases (Aguirre & Tabor, 2008), and vector-borne diseases (e.g. malaria, dengue, chikungunya) in particular (Harvell et al., 2002; McMichael, Woodruff, & Hales, 2006).

In the current study, we investigate the effect of temperature on the interaction between ranaviruses and their amphibian hosts, a host-pathogen system that offers the possibility of direct experimental manipulation, and a well characterized recent history of pathogen invasion into the UK (Price, Garner, Cunningham, Langton, & Nichols, 2016). Ranaviruses are large double-stranded DNA viruses (family *Iridoviridae*) that can be highly pathogenic to ectothermic vertebrates (Gray, Miller, & Hoverman, 2009; Price et al., 2014; Rosa et al., 2017). Ranavirus infections of amphibians are notifiable to the World Organization for Animal Health due to their potential to cause severe disease outbreaks as well as the risks of international spread through trade (Schloegel, Daszak, Cunningham, Speare, & Hill, 2010; Schloegel et al., 2009). Ranavirus growth and virulence can be affected by temperature (Ariel et al., 2009; Bayley, Hill, & Feist, 2013; Brand et al., 2016; Rojas, Richards, Jancovich, & Davidson, 2005) and environmental temperature is considered to be one possible explanation for observations of seasonality in outbreaks (Brunner, Storfer, Gray, & Hoverman, 2015). Indeed, incidents of ranavirosis in frogs in the USA were recently shown to be uncoupled from a pulse in transmission or the density of susceptible hosts, and instead were coincident with temperature increases and developmental changes in frog larvae (Hall, Goldberg, Brunner, & Crespi, 2018).

Ranaviruses are distributed globally but outbreaks of disease are extremely patchy - a pattern which is not yet understood. Some disease outbreaks have been shown to result from human translocations of ranavirus (Jancovich et al., 2005; Picco & Collins, 2008; Price et al., 2016), while other studies have found infections to be widespread at national scales without evidence for disease, which may reflect an historic association (Warne, LaBumbard, LaGrange, Vredenburg, & Catenazzi, 2016; Whitfield et al., 2013). The seasonal patterns and the observations of a temperature effect in laboratory studies raise the possibility that environmental conditions could drive invasion success and routes in cases where ranaviruses are undergoing range expansion as well as climate change being a driver of disease emergence in regions where the associations between viruses and hosts are historic and widespread.

In the UK, recurrent amphibian mass-mortality incidents caused by ranavirus have resulted in severe population declines of the common frog (*Rana temporaria*) (Teacher, Cunningham, & Garner, 2010). Genetic evidence supports multiple pathogen introductions into the UK whilst spatiotemporal models suggest that ranavirus spread rapidly, facilitated by translocations of unspecified infectious materials by people (Hyatt et al., 2000; Price et al., 2016). Disease outbreaks are strongly seasonal, peaking in the summer months and appearing to mostly affect adult animals, contrasting with other regions where larvae or metamorphic animals are the worst-affected age-classes (Brunner et al., 2015). However, the detectability of the main UK host, the common frog, is also strongly seasonal and there has been no previous attempts to explicitly control for host population density, host activity or observer effort in examining the periodicity of outbreaks (Cunningham, 2001; Cunningham et al., 1996; Teacher et al., 2010).

In this study, we investigated the role of temperature as a driver of disease outbreaks in common frogs infected with ranaviruses in the frog virus 3 (FV3) lineage through a combination of epidemiological modelling of a long-term study of disease in wild populations of UK common frogs (Cunningham, 2001; Price et al., 2016), *in vitro* experiments involving manipulation of the host-pathogen environment and similar *in vivo* experiments using natural hosts. Our aims were

to examine the role of temperature in shaping host-pathogen interactions to address whether it 1) has been a factor explaining the pattern of invasion which can be used to predict future changes in epidemiology under projections of climate change, and 2) can explain the observed seasonal patterns in disease occurrence and the contrast in affected host age-class compared to other temperate regions experiencing amphibian mortality incidents due to the same type of ranavirus.

2 Materials and Methods

2.1 Temperature as a predictor of frog mortality and incident severity

Temperature-dependence of ranavirus incidence: We used data from the Frog Mortality Project (FMP), a flagship citizen science project which has been collating reports of amphibian mortality incidents from members of the UK public for over twenty-five years (Price et al., 2016) to study disease occurrence in wild populations. The FMP dataset has been reliably filtered for incidents of ranavirosis previously (North, Hodgson, Price, & Griffiths, 2015; Price et al., 2016; Teacher et al., 2010). In this study, the same criteria as Cunningham (2001) and Price et al. (2016) (the presence of indicative signs of disease [‘ulceration’, ‘red spots on the body’, and ‘limb necrosis/loss of digits’; see Supporting information Appendix S1 and Figure S1] and a minimum of five dead animals) were used to create a binary variable describing the ranavirus status of each incident. The seasonal detectability of amphibians was controlled for indirectly through the inclusion of mortality incidents caused by factors other than ranavirosis as previously (Price et al., 2016). Data on the timing of the onset of mortality (available at the resolution of month only) and the incident location were used to download the monthly average of the daily maximum temperature for each incident from the Met Office UKCP09 dataset (Met Office, 2017b) (further details in Supporting information Appendix S2).

Factors affecting ranavirus incidence were investigated using a standard logistic regression model incorporating variables describing the environment (temperature), pond (volume, shading, and presence of marginal and floating vegetation), other aquatic vertebrate species present in addition to common frogs (toads, newts, fish) and geographic region (government office region) as predictors of ranavirus status (Model 1) fitted with the R function *glm2* (Marschner, 2011; R Core Team, 2017). To explore the relationship between temperature and the probability of a ranavirus-positive observation further, it was modelled as a sigmoid (logistic) transition between an upper and lower mean frequency. The four parameters of the curve (upper and lower limits, the location and the slope of the transition) were fitted using the *mle2* function in the R package, *bbmle* (Bolker, 2017; R Core Team, 2017). Starting values for the *mle2* algorithm were obtained for slope and location (intercept), using the *glm2* function with a bespoke link function for binomial data (Data S2). A confidence interval around the fitted line was generated using the delta method to compute variance of a function with the *emdbook* package (Bolker, 2008, 2016). The model was compared using Akaike’s Information Criterion (AIC) to a standard logistic curve (a simplified version of Model 1 with only temperature terms retained as predictors).

Effect of increased temperature on the severity of disease incidents: In order to analyze the effect of temperature on disease incident severity, which was calculated as the proportion of the population that died and therefore included the total number of dead animals, the dataset was filtered using only signs of disease as criteria for differentiating between incidents caused by ranavirus and those caused by other factors. Incidents were considered ranavirosis-consistent if any two of the three indicative signs of disease used above were reported. The total number of dead frogs and the number of surviving frogs (estimated by reporters) were combined as an estimate of the proportion of the population that died and used as the response variable term in a generalized linear model using the quasibinomial family to account for overdispersion in the data (Crawley, 2013). The average local maximum temperature for the month of onset of mortality and the predicted ranavirus status, as well as the interaction between main effects, were used as predictors of severity. This basic model was also extended by incorporating these same terms as well as other variables describing the pond environment (presence of other species and physical characteristics of ponds).

Temperature preceding ranavirus outbreaks with precise timestamps: We previously screened a UK amphibian and reptile tissue archive for ranavirus and returned a database of mortality incidents for which the presence of ranavirus was established using molecular diagnostic methods (Price et al., 2017). After filtering for incidents with a precise georeference (postcodes or grid references) and timestamp (date found, submitted, or examined if a post-mortem examination was conducted on receipt, which were each assumed to approximate closely to the day of death due to the rapid decomposition of amphibian carcasses), this dataset contained a total of 197 incidents, for which ranavirus had been detected from 31. All incidents were overlaid on the UK grid of 5 x 5 km squares and the maximum daily temperatures for the date matching the timestamp and the 50 days prior to the death(s) were extracted from plain text data files in the UKCP09 dataset (Met Office, 2017a), which were downloaded using the R package *Rcurl* (Temple Lang and CRAN team 2018; Price, 2018).

Ranavirus status was used as the response variable in a series of logistic regression analyses (generalized linear models using the binomial family and the logit link function) with the average maximum daily temperature in the week preceding the mortality incident or the number of consecutive days in the previous seven where temperature exceeded 16°C as predictors. These models were also run with region (Government Office Region) or latitude as an additional predictor to further control for any effect of spatial variation in temperature. As above, seasonal variation in the detectability of amphibians was controlled for by inclusion of mortality incidents caused by factors other than ranavirosis.

Effects of historic warming and seasonality: To check whether prior warming (over the time course of the dataset; 1991-2010) had altered the rate of ranavirus incidents and to assess seasonality in the data, we first decomposed the annual signal and trend across years in the time series of temperatures and rates of ranavirosis incidents. The mean of the average daily maximum temperature during the month of onset of mortality incidents from all reports in the FMP dataset (1991-2010) was calculated for each month with reports, as well as the numbers of reports that were consistent or otherwise with ranavirosis. Generalized Additive Mixed Models [GAMM; R function *gamm*; package *mgcv* (Wood, 2003, 2004, 2017)] were used to fit smooth

splines to both the within-year (seasonal; cyclic cubic regression spline) and across-year (cubic regression spline) patterns and autocorrelation structures (of order one) were used to model residual correlation within years. The number of ranavirosis incidents as a proportion of total reports was then modeled as a function of the seasonal trend (smoothed with a cubic regression spline as above), the trend in temperature across years (predicted using the GAMM above and smoothed with a cubic regression spline) and time using a Generalized Additive Model (GAM) and the binomial family with logit link function (function *gam*; package *mgcv*).

Model assumptions were verified by plotting residuals against fitted values and against each covariate in the model (function *gam.check*; package *mgcv*). Autocorrelation among residuals was assessed using the *acf* function in the *stats* R package (Venables & Ripley, 2002). Model predictions and residuals were extracted and visualized using functions in the R packages, *visreg* and *ggplot2* (Breheny & Burchett, 2013; Wickham, 2016). The predictive power of the model incorporating the across-year temperature trend was then assessed by comparison to a model containing the temporal trend in the rate of ranavirus incidents by dividing the dataset into two training and test sets (taking half and three-quarters of the data for training respectively). GAMs incorporating the smoothed seasonal trend and either a smoothed (across-year) time trend or the smoothed temperature trend were fitted using the training datasets and compared in terms of their ability to predict patterns in the test datasets.

2.2 Virus growth *in vitro*

To investigate the effect of temperature on viral growth, two UK isolates of FV3 [RUK11, isolated from the kidney of a diseased common frog that died with systemic hemorrhages in Middlesex in 1992, and RUK13, isolated from the skin of a diseased common frog that died with skin ulceration in Suffolk in 1995 (Cunningham, 2001)] were grown at a range of temperatures in two cell lines (epithelioma papulosum cyprini [EPC, derived from the fathead minnow fish, *Pimephales promelas* (Winton et al., 2010); ECACC 93120820] and the iguana heart reptile line [IgH2; ECACC 90030804]). Cells were grown on 96 well plates until more than 90% confluent and then inoculated with virus in a ten-fold dilution series ranging from an estimated multiplicity of infection of approximately 2×10^{-6} to 2×10^3 (five wells per dilution with an additional one well per dilution receiving a sham exposure of cell culture media only as a negative control). Titers of viral isolate stocks were equalized by reference to qPCR scores [following Leung et al. (2017)]. Plates were incubated at six temperatures (10, 14, 18, 22, 26, and 30°C) and monitored daily for cytopathic effect (plaques in the cell layer). After six days, plates were scored for viral growth by counting the number of replicates at each dilution where cytopathic effect was evident and calculating the Tissue Culture 50% Infective Dose (TCID50) using the method of Reed & Muench (1938). The effects of temperature and cell line on viral growth (the mean titers of the two isolates of each type) were analyzed using a linear model in R.

2.3 *In vivo* assessment of effect of temperature on virulence

To investigate whether an effect of temperature on *in vitro* viral growth was reflected in altered virulence *in vivo*, 60 overwintered common frog metamorphs (*R. temporaria*) were randomly allocated to one of six treatments (10 animals per treatment): three exposure treatments (sham,

RUK11, RUK13) crossed with two temperatures (20°C [“low”] and 27°C [“high”]). Temperature was maintained by placing five individually-housed frogs selected at random from each of the three exposure treatments into one of four climate-controlled chambers constructed from polystyrene boxes - two held at 20°C and two held at 27°C (Figure S2; see Supporting information Appendix S3 for a comprehensive description of the set up). RUK13 was used at a titer of 1.58×10^5 TCID₅₀ mL⁻¹ and RUK11 at a titer of 1.58×10^7 TCID₅₀ mL⁻¹ (see Supporting information Appendix S3 for details of preparation of inocula and exposure methods).

Individuals reaching endpoints (either gross signs of hemorrhaging or ulceration; Figure S1) and all surviving individuals at the end of the experiment (day eight post-exposure) were euthanized following a schedule 1 method for amphibians. The number of hours post-exposure that animals were found dead or at endpoints was recorded for survival analysis. Survival data were analyzed by fitting a mixed-effects Cox proportional hazards regression model in R using the *coxme* package (Therneau, 2018) with exposure and temperature as fixed effects and the climate-controlled chamber as a random effect to account for pseudo replication due to placement of individual experimental units inside the four climate-controlled chambers. Model coefficients were visualized using *forestplot* (Gordon & Lumley, 2017).

A second *in vivo* experiment was performed in the same host species, using one ranavirus isolate (RUK13) and a single temperature (20°C), but two exposure doses (“low” and “high”; detailed methods in Supporting Information Appendix S1). The effects of temperature and dose on disease progression and survival were compared.

2.4 Effect of projected climate change on timing of ranavirus outbreaks

Baseline temperature data were generated by calculating the mean daily maximum temperature for each calendar month for the period 1991-2010 (the same period covered by our dataset of reported frog mortality incidents) for each 5 x 5 km grid square used by the Met Office UKCP09 project (Met Office, 2017b). Probabilities of mean daily maximum temperature exceeding 16°C for calendar months during the period 2070-2099 under the Intergovernmental Panel on Climate Change (IPCC) B1 (“low”), A1B (“medium”) and A1FI (“high”) future emissions scenarios (IPCC, 2000) were downloaded for a grid of 25 x 25km squares at UK scale using the UK Climate Projections user interface (Jenkins et al., 2009).

3 Results

3.1 Effect of temperature on disease occurrence and severity in the wild

The finalized FMP dataset used in this study contained 4385 unique records, of which 1497 were classed as ranavirosis-consistent. The logistic model (Model 1) revealed a highly significant, non-linear effect of temperature on the proportion of ranavirosis-consistent incidents observed (Table S1). The minimal adequate model also retained newts, fish and shading: the presence of either type of animal in ponds increased the proportion of ranavirosis-consistent incidents observed whilst shading reduced this proportion (Table S1). To explore the relationship with temperature further, a model with a transition between an upper and lower frequency was fitted (Model 2), which significantly improved the fit to the data compared to a simplified version of Model 1 comprising only the terms describing the non-linear relationship with temperature (AIC scores were 5565 and 5576 respectively). Model 2 shows a step-change: below approximately 16°C, 25.1% of incidents were ranavirosis-consistent, rising to 38.5% after the temperature threshold was crossed (Figure 1a). The difference between incidents that were ranavirosis-consistent and the remainder ('non-ranavirus') is also apparent in the distribution of temperature records: the non-ranavirus category being strongly bimodal with peaks at both low and high temperature, whereas most of the ranavirosis-consistent incidents were reported at higher temperatures with few outliers at much lower temperatures (Supporting information Figure S3).

Temperature was again a highly significant predictor of ranavirus status when the records with precise timestamps and confirmed ranavirus-positive status were analyzed. This more precise information about timing enabled a fine-scale examination of the effect of temperature in the days preceding incidents. The average temperature in the seven days preceding incidents was a significant predictor of ranavirus status ($p = 1.18 \times 10^{-4}$; residual deviance of model = 154.2 on 195 degrees of freedom; Figure 1b), with each 1°C increase in temperature increasing the odds that incidents were caused by ranavirus by 20%. The temperature threshold where the proportion of ranavirus incidents increased sharply in the analysis of the full FMP dataset was approximately 16°C. A second model - with the number of consecutive days where the daily maximum temperature in the week preceding incidents exceeded 16°C as a predictor - also indicated that warmer temperatures were a good predictor of ranavirus status ($p = 2.27 \times 10^{-5}$; residual deviance of model = 151 on 195 degrees of freedom; Figure 1c): each additional warm day raised the odds that incidents were caused by ranavirus by 33%. The 16°C threshold model had a slightly lower AIC score than the model using the average temperature as a predictor (155 compared to 158).

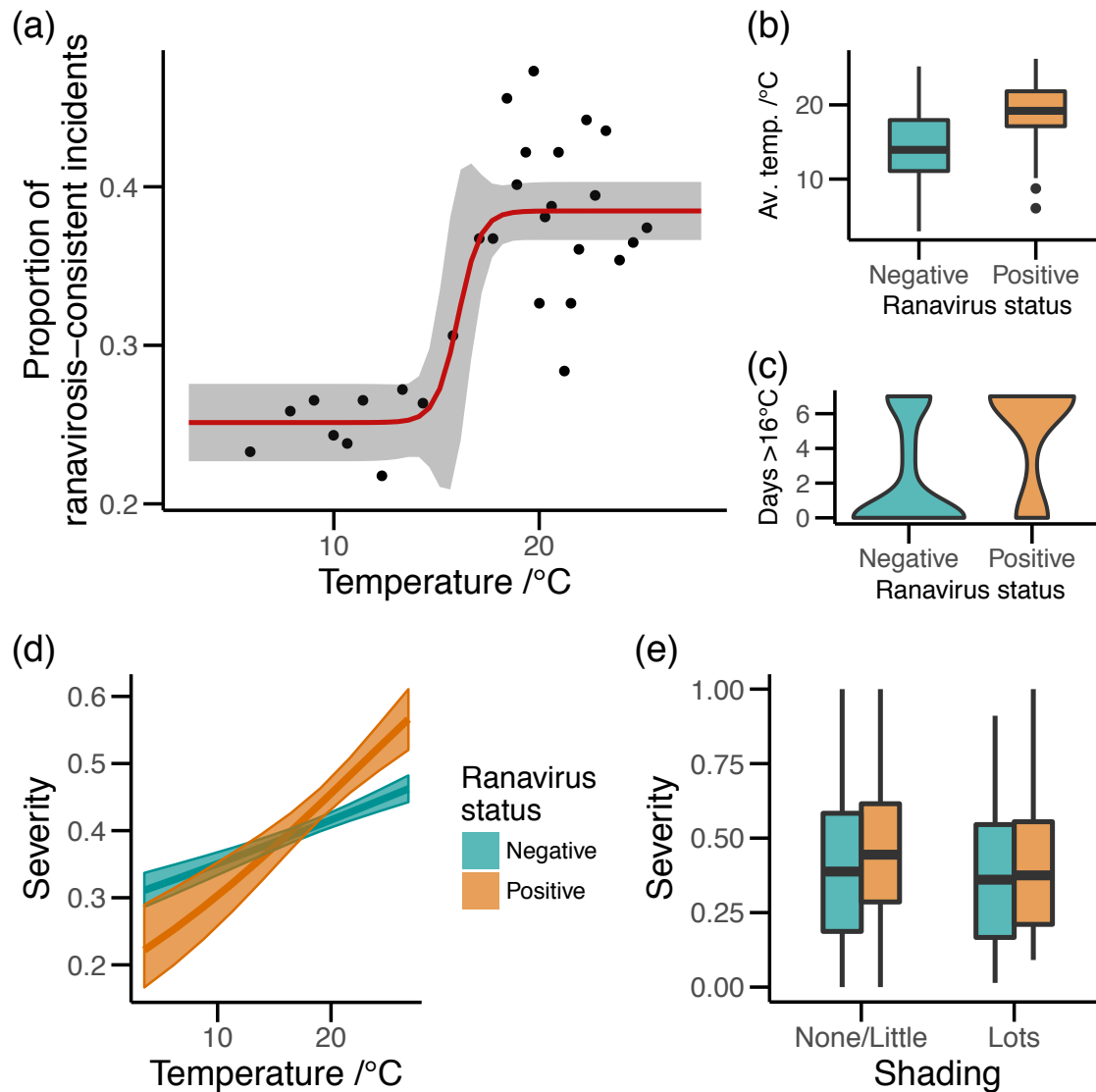


Figure 1. Warm temperatures increased the frequency and severity of incidents of ranavirosis involving wild common frog populations of the United Kingdom. (a) The effect of temperature on the proportion of citizen science reports of frog mortality that were classified as ranavirosis-consistent. The line represents the fitted maximum likelihood model of a logistic transition between a lower and upper frequency. The shaded area around the line represents the 95% confidence interval, calculated using the delta method. Points represent the observed data, grouped in windows containing equal numbers of records. (b-c) Temperature in the week preceding frog mortality incidents confirmed by molecular methods predicted ranavirus status (“Positive” or “Negative”). (b) Average daily maximum temperature in the seven days preceding incidents by ranavirus status. (c) The number of days in the week preceding mortality incidents where the daily maximum temperature exceeded 16°C by ranavirus status. (d) The severity of frog mortality incidents (estimated proportion of population that died) was consistently greater at higher temperatures, particularly in the case of ranavirosis-consistent incidents (orange). The

plot shows fitted lines (and 95% confidence intervals) from a generalized linear model (quasibinomial regression) of severity as a function of ranavirus status and temperature (average daily maximum temperature for the month of onset of mortality incidents). (e) Large amounts of shading around ponds reduced the severity of ranavirosis-consistent mortality incidents (orange) but had no effect on other incidents (green). In panels (b) and (e), boxplots represent lower quartile, median, upper quartile and interquartile range (upper quartile - lower quartile; central 50% of the data); whiskers extend to the most extreme data point which is no more than 1.5 times the interquartile range from the box; outliers shown as individual points where relevant.

The FMP database contains data on the severity of outbreaks (the estimated proportion of the frog population that died) for the years 1991-2000. After removing records with missing values, we produced a dataset for investigating severity that contained 2667 records, of which 427 incidents were classified as ranavirosis-consistent. In a simple logistic model of severity with ranavirus status, average daily maximum temperature for the month of onset of mortality, and their interaction as predictors, all three terms were significant predictors and retained in the minimal adequate model. Temperature explained the most deviance with each 1°C increase in temperature leading to a 2.8% increase in the proportion of the population that died ($p = 1.61 \times 10^{-12}$). There was a significant interaction between the two main effects as a consequence of the different effect of ranavirus status on the relationship between temperature and the proportion dead ($p = 0.001$): at low temperatures, the severity of ranavirosis-consistent incidents was slightly lower than for other types of incident but at higher temperatures it was the ranavirosis-consistent incidents that were more severe (Figure 1d; Supporting information Figure S4). Attempting to incorporate time (the year that mortality incidents began) did not result in an extension of the minimal adequate model.

The effects of other covariates previously identified as having an influence on the occurrence or severity of ranavirosis in UK common frogs (North et al., 2015) were also explored using a more complex model containing ranavirus status, temperature, log-transformed pond volume, the interactions of the three, shading around ponds, the amount of both the marginal and floating vegetation (“none/little” or “lots”), the presence of toads, the presence of newts, the presence of fish, and the region. After model simplification, the minimal adequate model retained all three terms from the simple model but there were also significant effects of the presence of toads, the presence of fish, shading, pond volume (non-linear), marginal vegetation and region (Table S2). Toads reduced the severity of mortality incidents whilst the presence of fish increased severity (Supporting information Figure S5) as found previously (North et al., 2015). Shading and marginal vegetation decreased the severity of incidents. Notwithstanding, and irrespective of which covariate was considered, the effect of increasing temperature increased the severity of disease. This is perhaps best illustrated by the effects of pond shading, where increasing the amount of shading (and, presumably, decreasing the maximum temperatures that frogs would have been exposed to) was associated with reduced severity of ranavirosis and a reduction in the disparity in the severity of incidents between ranavirosis-consistent and non-ranavirus incidents (Figure 1e).

3.2 Historic climate change and seasonality

Temperatures in our study region varied across years, showing a warming trend that peaked in 2002 before cooling up to 2010, as well as showing marked seasonality (GAM component of GAMM: $R^2 = 0.93$; Table S3). The proportion of ranavirosis incidents followed a remarkably similar pattern, in terms of the strong seasonality and the trend across years (GAM component of GAMM: $R^2 = 0.36$; Table S4; Figure 2a). The across-year temperature trend was a highly significant predictor of the rate of ranavirus incidents (GAM: effective degrees of freedom (edf) of smoothed term = 6.65, $p = 4.61 \times 10^{-26}$), explaining the pattern of increasing rates which peaked in 2001, with the full model explaining approximately half the deviance ($R^2 = 0.48$; Figure 2; Table S5). Model validation indicated conformity to assumptions. The temperature model, when trained on subsets of the data, predicted trends in test datasets effectively in contrast to models incorporating only the across-year trend in ranavirosis rates, which showed no predictive power (Supporting information Figure S6). Temperature showed a strong seasonal pattern which was reflected in the pattern of ranavirosis incidents where seasonality was also marked (GAM: edf = 7.65, $p = 2.45 \times 10^{-13}$; Figure 2c), raising the possibility that another seasonal factor - correlated with temperature - could have driven ranavirus outbreaks. However, the association between temperature and the rate of ranavirus incidents across years suggests that temperature is driving the rate of disease incidents in the long term and may also drive the seasonality, since both a correlated seasonal factor and a correlated non-seasonal factor acting across years would otherwise be required to explain the observed patterns. Overall, after accounting for the effect of temperature, there was a small but significant decrease in the proportion of ranavirosis incidents between 1991 and 2010 (GAM: coefficient = -0.00315 [unit of time was months], $p = 2.31 \times 10^{-5}$; Figure 2d).

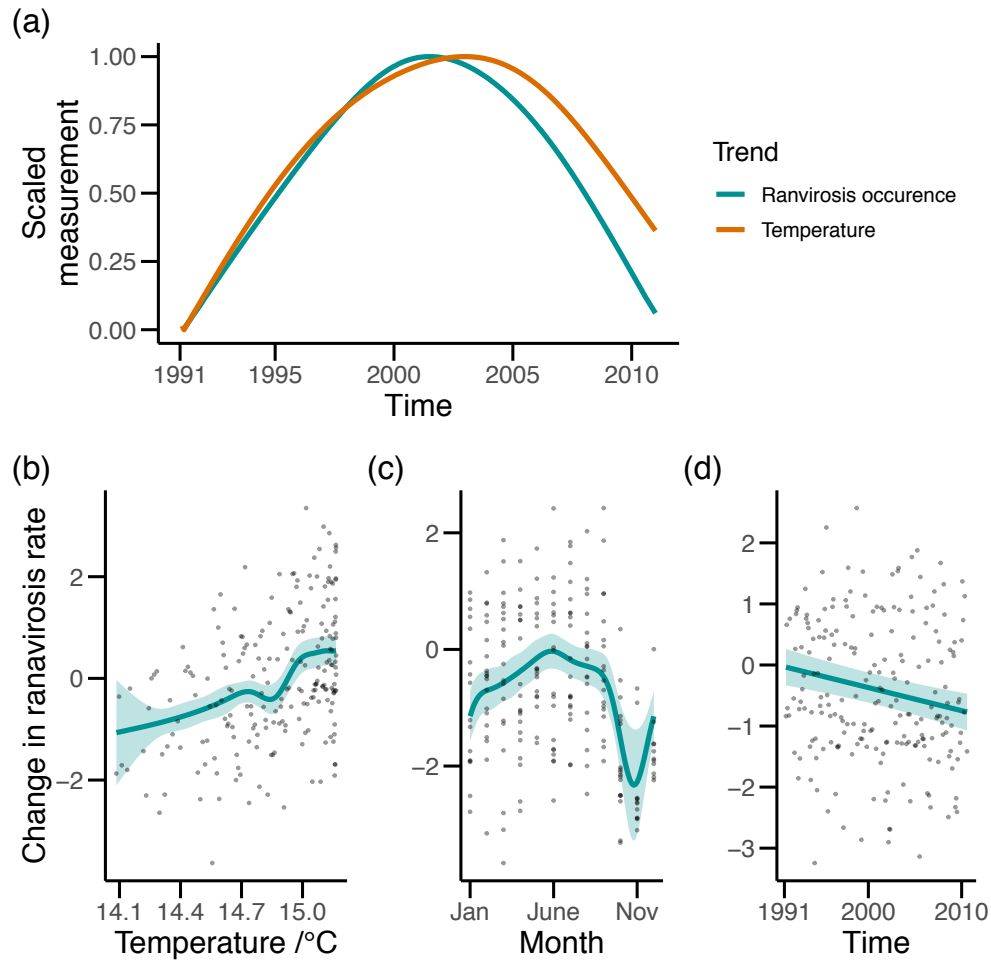


Figure 2. Effect of historic climate and seasonality on rate of ranavirosis incidents. (a) comparison of smoothed trends in temperature and rates of ranavirosis incidents (on standardized scale [0-1]) over period of dataset (1991-2010). (b-d) Effect of predictors of the rate of ranavirosis incidents (from generalized additive model) against residuals: Smoothed change in rate of ranavirus incidents with temperature (b), smoothed seasonal change in rate of ranavirosis incidents (c), and change in rate of ranavirosis incidents over time (d). Shaded areas represent 95% confidence intervals.

3.3 *In vitro* assessment of viral growth rates and *in vivo* tests of virulence

We examined *in vitro* viral growth using two UK isolates of FV3 (RUK11 and RUK13; see methods for detailed descriptions of isolates) and two cell lines. Each isolate was incubated at a range of temperatures up to 30°C with each cell line, but regardless of cell line or isolate, increasing temperature resulted in exponentially increased rates of plaque formation (Figure 3). A linear model of log viral titers against temperature, cell line and their interaction revealed significant

effects of temperature (coefficient = 1.23, $p < 1e^{-30}$) and host cell line (IgH2 compared to EPC, coef = -8.64 , $p < 1e^{-30}$) but no interaction (analysis of variance, comparing model with interaction term to a model with main effects only: $F_{df=1} = 0.0047$, $p = 0.95$), indicating the overall effect of temperature was independent of the host environment.

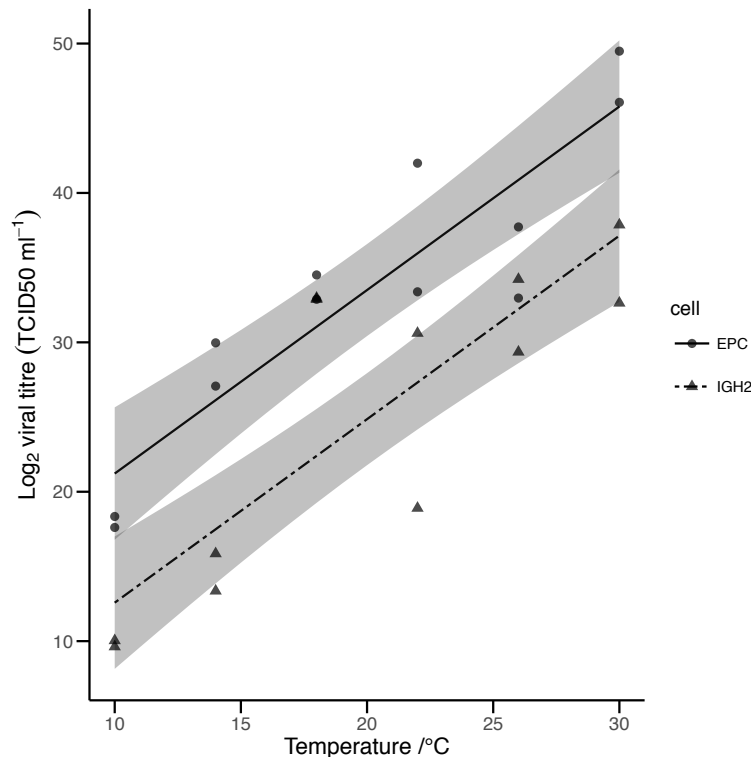


Figure 3. Effect of environmental temperature on growth of UK Frog virus 3 (FV3) *in vitro*. Observed data (points) and predictions from linear model (lines with 95% confidence interval shaded) of FV3 growth at a range of environmental temperatures in fish (EPC; solid line) and reptile (IgH2; dashed line) cells. Growth was measured using the TCID50 method and is shown on a log scale. An increase in temperature of 1°C results in more than a doubling of viral growth (2.34 times).

The effect of temperature on the response of common frogs to viral exposure was assessed in order to validate results from cell culture models *in vivo*. Temperature was a highly significant predictor of survival: 20 of 60 animals died or were euthanized on reaching humane endpoints, of which 14 were from high temperature treatments and six were from low temperature treatments (Figure 4a). Overall there was a 5.33 times higher risk of death in the high temperature treatments ($p = 0.005$; Figure 4b). Titers of viral inoculates were not equalized between isolates. All individuals exposed to RUK11 (at a high dose) and maintained at high temperature died or reached endpoint by the eighth day post-exposure compared to six of ten individuals maintained at low temperature (Figure 4a). Of the animals exposed to RUK13 (at a

relatively low dose compared to RUK11), three individuals died or reached endpoint in the high temperature treatment compared with none at the low temperature. There was also a significant effect of exposure treatment: the expected hazard of animals exposed to RUK11 was 41.6 times higher than animals receiving a sham exposure ($p = 0.0004$). These results are largely in line with a study examining survival of common frog tadpoles exposed to a North American isolate of FV3, which showed that mortality was increased at 20°C compared to 15°C (Bayley et al., 2013).

The second *in vivo* experiment examining the effect of dose on disease outcome and progression in juvenile common frogs (Supporting Information Appendix S1) complements the findings of the *in vivo* temperature experiment. The dose experiment suggests that a viral load threshold exists which must be crossed before gross signs of disease develop. We found the outcome and presentation of disease as well as the viral quantity in tissues at death to be largely independent of dose: all animals exposed to either low or high viral doses died, presented with the same set of signs (Figure S1 & S7), and had similar quantities of virus in their tissues at death (Figure S8a). However, the onset and progression of disease was delayed at low dose (the development of disease and death both occurred significantly later; Figure S7) reflecting the lower viral loads of individuals (measured by swabbing animals after infection; Figure S8b) given this treatment. Also, viral loads of dead individuals were greater than those of live individuals (both when repeated-measures from the same individuals were compared [Figure S8c] and when individuals that were euthanized part-way through the experiment were compared to those that died [Figure S8d]). These results all suggest that elevated viral loads lead to the onset of disease and that the viral capacity to cross a threshold concentration is a more important determinant of whether disease develops than the initial dose. It seems likely that higher temperature and higher initial dose each serve as ways to reach this putative threshold for disease sooner - either through more rapid viral growth or a greater initial intensity of infection respectively - and explain the delays and/or reductions in observations of severe outcomes in the other respective treatments (low temperature or low dose).

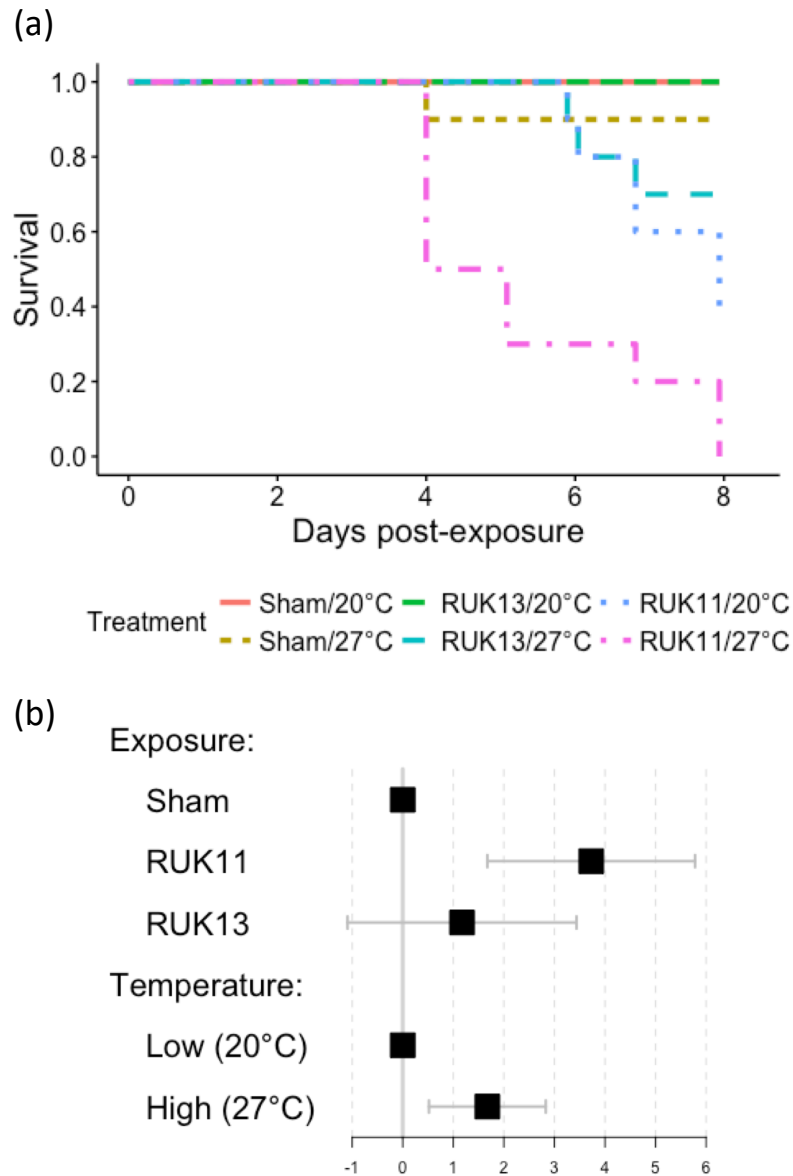


Figure 4. Effect of temperature and ranavirus exposure on survival of common frogs. The proportion of surviving animals through time plotted for each of six treatments (n=10 frogs per treatment); three exposure treatments (Sham, RUK11, RUK13) at each of two temperatures (20°C [“low”] or 27°C [“high”]). (a) Kaplan-Meier survival plot. (b) Forest plot of coefficients (\pm standard error) from a mixed effects Cox Proportional Hazards model of survival in response to exposure (“Sham” as the reference level) and temperature (“Low” as the reference level) treatments.

3.4 Impact of future climate on timing of outbreaks

The UK climate is expected to warm considerably over the remainder of the century (Chen & Tung, 2018; Jenkins et al., 2009). A warmer climate would expand the geographic area where environmental conditions are likely to be suitable for severe incidents of ranavirosis (average monthly maximum daily temperatures exceeding 16°C). For example, the suitable geographic area for ranavirosis occurrence in May is projected to increase by 134% by 2070 under a high emissions scenario compared to the historic baseline and by 84% under a low emissions scenario (Figure 5). The projected changes in UK temperatures also will extend the duration of the “disease season”, creating favorable conditions for disease during the spring and autumn as well as in the summer. Temperatures are likely to become more suitable for severe outbreaks in a large part of England in April under a high emissions scenario and in October under any of the range of IPCC scenarios investigated. These are months that we expect to have experienced limited incidence and severity of ranavirosis previously, but temperatures are projected to change to such a degree that this limitation will be removed for large areas of the UK (Figure 5).

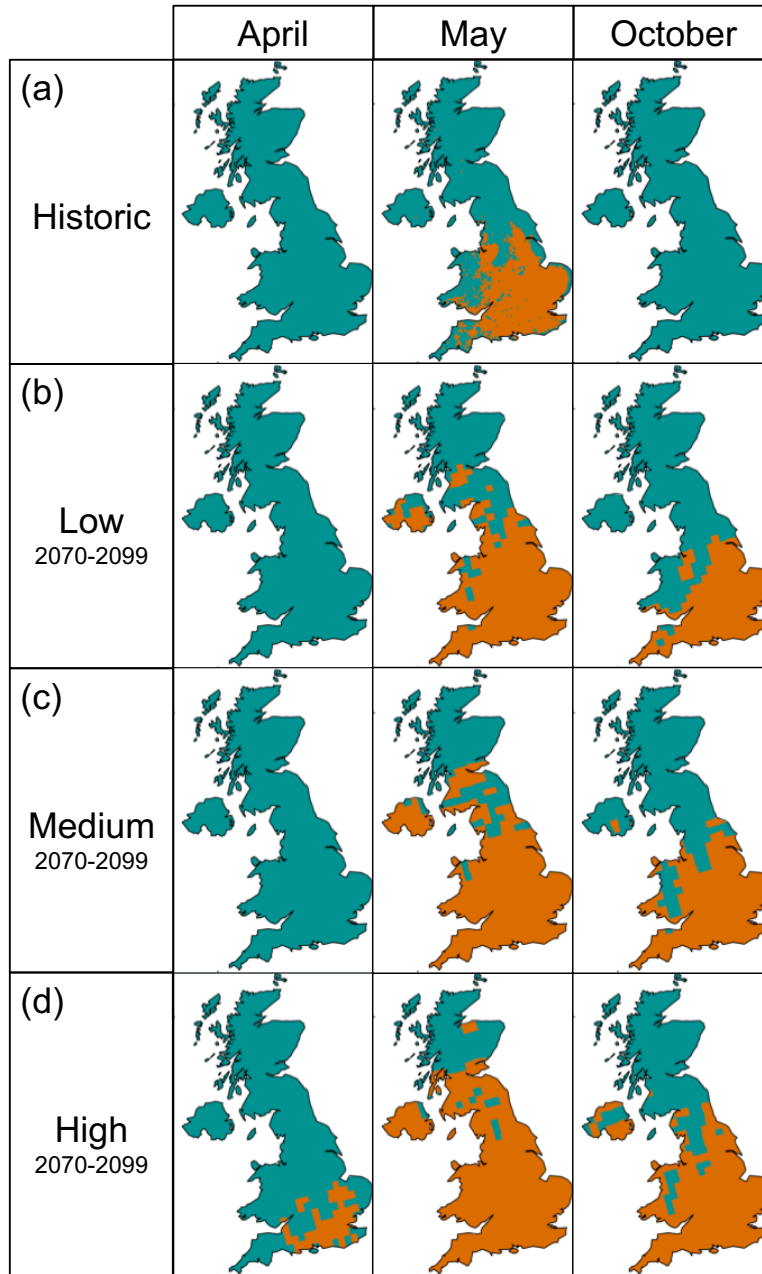


Figure 5. Projected shifts in geographic extent and the temporal window of ambient temperatures high enough to increase the risk and severity of ranavirosis incidents in the UK under different emissions scenarios. (a) Past climate - orange regions are those where the average monthly temperatures (daily maximum temperature) in the UK (5 x 5 km grid squares) exceeded 16°C for the period 1991-2010. (b-d) Future climate - orange regions are those where the projected average monthly daily maximum temperatures in the UK (25 x 25 km grid squares) have a greater than 50% probability of exceeding 16°C under a range of future emissions scenarios for the period 2070-2099: (b) Intergovernmental Panel on Climate Change (IPCC) scenario B1 (low), (c) scenario A1B (medium) and (d) scenario A1FI (high).

4 Discussion

By combining the results of laboratory experiments with the analysis of two types of epidemiological dataset relating to disease outbreaks in wild amphibian populations, we have revealed a pattern of remarkably consistent evidence supporting a substantial effect of temperature on ranavirus disease dynamics. Temperature predicts both the incidence and severity of disease outbreaks through higher temperatures increasing viral growth which, in turn, manifests as an increased rate of disease occurrence in both experimental and wild frog populations. The use of *in vitro* and *in vivo* studies in combination with modeling of field datasets serves as a ‘triangulation’ process (Plowright, Sokolow, Gorman, Daszak, & Foley, 2008) and strongly suggests a causal link between temperature and disease occurrence in this system. Temperature is, of course, correlated with a multitude of other factors that may be considered alternative or additional drivers of disease outbreaks, but our findings of an association between temperature and disease incidents at two different timescales (within and across years) and the triangulation process both support the conclusion that temperature has a causal effect on ranavirus disease occurrence.

Our results are consistent with an historic effect of climate on the rate and timing of ranavirus incidents and suggest that the invasiveness of this introduced pathogen may have been facilitated, but also restricted, by the suitability of local climate. We have previously shown that ranavirus was introduced to the UK and spread rapidly in England (Price et al. 2016). The small overall decrease in the rate of disease incidents observed over the twenty years of data analyzed (Figure 2d) may be due to fewer opportunities for spread arising from a more complete colonization of the suitable range or might be credited to attempts to limit the risk of translocations through advice disseminated in the media (e.g. BBC, 2008; Price et al., 2016). Alternatively, the severe impacts of recurrent ranavirus outbreaks - previously shown to have caused declines of common frogs in South-East England (Teacher et al., 2010) - may have led to the extirpation of populations or reduced their size/density to the point where transmission no longer occurred, or disease outbreaks were no longer detectable.

Climate projections show how climate change will likely play a role in shaping future ranavirus disease dynamics in UK common frogs, altering both the geographic extent and the length of the temporal window of heightened disease risk and severity. The potential impacts of climate warming on disease ecology, therefore, could have critical ramifications for the continued survival of amphibian populations across the UK. Although it is challenging to detect disease and mortality in larval amphibians, all evidence points to adult common frogs as the major life history stage and species affected by ranavirosis in the wild in the UK (Cunningham, 2001; Duffus, 2009; Price et al., 2017). This observation is intriguing since larval forms are usually more affected by FV3 elsewhere in the world and common frog larvae have been shown to be highly susceptible to wild-type ranaviruses in the laboratory (Duffus, Nichols, & Garner, 2013; Duffus, Nichols, & Garner, 2014; Gray et al., 2009).

Our findings suggest that temperature could be an important determinant of the partitioning of disease among life-history stages, with common frog larvae metamorphosing before pond water reaches a temperature high enough to trigger outbreaks of ranavirosis. This situation could be

altered as the climate warms and the disease season is lengthened. Shifting the timing of frog disease outbreaks will alter the life history stages at risk. If common frog tadpoles become affected, the abundance of susceptible hosts will be increased with concomitant impacts on the ranavirus basic reproductive number (R_0) (Altizer et al., 2006); i.e. the dynamics of outbreaks will be fundamentally changed, making predictions of their impacts more challenging. Whilst the breeding phenology of some UK amphibian species has altered in response to climatic changes, common frog breeding may not respond to increasing absolute temperatures (Beebee, 1995), so any compensatory change in host behavior may be negligible. Common frog populations in regions where temperatures become suitable for severe disease outbreaks during the larval stage might experience reduced recruitment and a subsequent reduction in their capacity to persist in the presence of infection. Additionally, altering the timing of outbreaks could create opportunities for host jumps if other potential host species have increased contact with (high levels of) virus (Hoberg & Brooks, 2015).

The effects of temperature seem to act on the virus predominantly rather than on the host, as evidenced by the cell culture experiments which limit host factors that might be altered by temperature and therefore attribute changes in ranavirus growth to the pathogen itself. Increased virulence at higher temperatures is unlikely to be counteracted by any negative effects of higher temperature on ranavirus, as viral particles are persistent in the environment and are highly tolerant of exposure to extreme temperatures despite failing to replicate at temperatures above 30°C (Cunningham, 2001; La Fauce, Ariel, Munns, Rush, & Owens, 2012). Nor does it seem likely that more effective host immune responses that may accompany increased temperatures will counteract the effects on ranavirus if the capacity of ranaviruses to successfully evade common frog immune responses that we have shown previously operates across a range of temperatures (Price et al., 2015).

Nevertheless, our laboratory findings that virulence is reduced at lower temperatures, that frogs might be better able to manage infections at lower temperatures, together with field records showing a mitigating effect of shading, pond volume and vegetation (which might also be due to lowered temperatures of frogs) on incidence and severity of ranavirosis, point to possible steps for mitigation. Thermoregulatory behavior leading to an increase in body temperature above normal range (“behavioral fever”) was shown to reduce the odds of infection with chytrid fungus in Panamanian golden frogs (Richards-Zawacki, 2010) and is known to be important for disease mitigation in other ectotherms (Elliot, Blanford, & Thomas, 2002). Whether “behavioral cooling” also serves as an amphibian strategy for managing infections remains to be elucidated and staying cool can be a more difficult challenge than getting warm for many ectotherms (Kearney, Shine, & Porter, 2009). However, the provision of suitable opportunities for behavioral regulation of body temperature in the form of shading, log piles and larger ponds might help to manage the severity of future outbreaks.

Ranavirosis has had a major impact on common frog populations in south-east England (Teacher et al., 2010) and the current study suggests that these impacts may become greater and more widespread (in the UK and elsewhere) if future climate change projections are realized. Our results - as well as the predictions that follow from them - are strengthened through the use of a model system that allows us to investigate possible drivers of field epidemiology using laboratory

experiments, both at the cellular and whole animal levels. Together, our results present a clear case of the environment modulating an important host-pathogen interaction. Few previous studies have convincingly shown how climate change affects disease emergence in wild animal populations, but we have been able to demonstrate a historic impact of warming in the wild and then tease apart relationships between the environment, host and pathogen in the laboratory. The results highlight how species with complex life cycles might undergo sudden shifts in the level of threat posed by an infectious disease if gradual changes to the climate result in greater exposure/susceptibility of alternative life history stages when a favorable environment had previously buffered them against the most severe impacts of disease.

Acknowledgements

We thank Rob Knell for help and advice about climate-chamber construction. This work was funded by NERC grants NE/M000338/1, NE/M000591/1 & NE/M00080X/1. All *in vivo* experimental procedures and husbandry methods were approved by the ZSL Ethics Committee before any work was undertaken and procedures were performed under UK Home Office licenses P8897246A & 80/2214. The authors declare that there are no conflicts of interest. All data and code required to reproduce the analyses and figures in this article are included in the Supporting information files (Data S1 and Data S2). We thank Froglife (Registered Charity No. 1093372 in England and Wales) for careful administration and promotion of the Frog Mortality Project.

Author contributions

SJP & AAC designed the *in vitro* experiments, the lab work was performed by CO & WL, and analysis was performed by SJP with help from CO. SJP designed the *in vivo* experiments with help from TWJG & RAN, the lab work was performed by SJP, WL & CS, and the results were analyzed by SJP. AAC oversaw collection of the epidemiological datasets, SJP, RP, TWJG, RAN & FB planned the analyses which were conducted by SJP and RAN. SJP and RP planned the climate change projections, which were performed by SJP. SJP & FB wrote the first draft of the manuscript which was edited by all authors.

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